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(54) Title: α -KETOAMIDE INHIBITORS OF 20S PROTEASOME

$$X_2$$
 N
 R_1
 R_1
 R_1

(57) Abstract

 α -ketoamide compounds useful for treating disorders mediated by 20S proteasome in mammals having structure (I).

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SPECIFICATION

TITLE: α-Ketoamide Inhibitors of 20S Proteasome

Background of the Invention:

The multicatalytic proteinase or the proteasome is a highly conserved cellular structure that is responsible for the ATP-dependent proteolysis of most cellular proteins (Coux, O., Tanaka, K. and Goldberg, A. 1996 Ann. Rev. Biochem. 65, 801-847). The 20S proteasome contains the catalytic core of the complex and has been crystallized from the archaebacteria Thermoplasma acidophilum (Lowe, J., Stock, D., Jap, B., Zwicki, P., Bauminster, W. and Huber, R. 1995 Science 268, 533-539) and from the yeast Saccharomyces cerevisiae (Groll, M., Ditzel, L., Lowe, J., Stock, D., Bochtler, M., Bartunik, HD and Huber, R. 1997 Nature 386, 463-471). Unlike the archaebacterial proteasome that primarily exhibits chymotrypsinlike proteolytic activity (Dahlmann, B., Kopp, F., Kuehn, L., Niedel, B., Pfeifer, G. 1989 FEBSLett. 251, 125-131; Seemuller, E., Lupas, A., Zuw, F., Zwickl, P and Baumeister, W. FEBS Lett. 359, 173, (1995) the eukaryotic proteasome contains at least five identifiable proteolytic activities. Three of these activities are similar in specificity to chymotrypsin, trypsin and peptidylglutamyl peptidase. The two other activities described exhibit a preference for cleavage of peptide bonds on the carboxyl side of branched chain amino acids (BrAAP) and toward peptide bonds between short chain neutral amino acids (SnAAP) (Orlowski, M. 1990 Biochemistry 29, 10289-10297).

Although the 20S proteasome contains the proteolytic core, it cannot degrade proteins in vivo unless it is complexed with a 19S cap, at either end of its structure, which itself contains multiple ATPase activities. This larger structure is known as the 26S proteasome and will rapidly degrade proteins that have been targeted for degradation by the addition of

multiple molecules of the 8.5-kDa polypeptide, ubiquitin (reviewed in Coux, O., Tanaka, K. ard Goldberg, A. 1996 Ann. Rev. Biochem. 65, 801-847).

A large number of substrate-derived functionalities have been used as potential serineand thiol protease inhibitors. Several of these motifs have been described as inhibitors to the proteasome. These include the peptide aldehydes (Vinitsky, A., Michaud, C., Powers, J. and Orlowski, M. 1992 Biochemistry 31, 9421-9428; Tsubuki, S., Hiroshi, K., Saito, Y., Miyashita, N., Inomata, M., and Kawashima, S. 1993 Biochem. Biophys. Res. Commun. 196,1195-1201; Rock, K,I., Gramm, C., Rothstein, L., Clark, K., Stein, R., Dick, L., Hwang, D. and Goldberg, A.L. (1994) Cell 78, 761-771) N-acetyl-L-leucinyl-L-leucinyl-L-norleucinal (ALLN) and N-acetyl-L-leucinyl-leucinyl-methional (LLM) with the most potent inhibitor of this type being N-carbobenzoxyl-l-L-leucinyl-L-leucinyl-L-norvalinal (MG115). Other reports have described a series of dipeptide inhibitors that have IC_{so} values in the 10 to 100 nM range (Iqbal, M., Chatterjee S., Kauer, J.C., Das, M., Messina, P., Freed, B., Biazzo, W and Siman, R. 1995 I-Med.Chem. 38, 2276-2277). A series of α-ketocarbonyl and boronic ester derived dipeptides (Iqbai, M., Chatterjee, S., Kauer, J.C., Mallamo, J.P., Messina, P.A., Reiboldt, A. and Siman, R. 1996 Bioorg. Med-Chem. Lett 6, 287-290) and epoxyketones (Spattenstein, A., Leban, JJ., Huang, J.J., Reinhardt, K.R., Viveros, O.H., Sigafoos, J. and Crouch, R. 1996 Tet. Lett. 37, 1434-1346) have also been described that are potent inhibitors of the proteasome.

A different compound that exhibits specificity in inhibiting proteasome activity is Lactacystin (Fenteany, G., Standaert, R.F., Lane, W.S., Choi, S., Corey, E.J. and Schreiber, S.L. 1995 Science 268, 726-731) which is a Streptomyces metabolite. This molecule was originally discovered for its ability to induce neurite outgrowth in a neuroblastoma cell line (Omura, S., Matsuzaki, K., Fujimoto, T., Kosuge, K., Furuya, T., Fujita, S. and Nakagawa, A. 1991 J.Antibiot. 44, 117-118) and later was shown to inhibit the proliferation of several cell

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types (Fenteany, G., Standaert, R.F., Reichard, G.A., Corey, E.J. and Schreiber, S.L.1994 Proc.Nat'l. Acad.Sci.USA 91, 3358-3362).

It is now well established that the proteasome is a major extralysosomal proteolytic system involved in the proteolytic pathways essential for diverse cellular functions such as cell division, antigen processing and the degradation of short-lived regulatory proteins such as oncogene products, cyclins and transcription factors (Ciechanover, A. (1994) Cell 79, 13-21;Palombell,V.J., Rando, O.J., Goldberg, A.L. and Maniatis, T. 1994 Cell 78, 773-785). For example, the active form of NF-kB is a heterodimer consisting of a p65 and a p50 subunit. The latter is present in the cytosol as an inactive precursor (pl05). The proteolytic processing of p105 to generate p50 occurs via the ubiquitin-proteasome pathway. Additionally, processed p50 and p65 are maintained in the cytosol as an inactive complex bound to the inhibitory protein IkB. Inflammatory stimuli such as LPS activate NF-kB by initiating the signalling pathway which leads to the degradation of IkB. These signals also stimulate the processing of p105 into p50. Thus two proteolytic events, both governed by the ubiquitin-proteasome pathway, are required for signal induced activation of NF-kB.

The observation that ubiquitin-mediated proteasomal proteolysis plays a critical role in the activation of NF-kB could be exploited clinically by the use of inhibitors directed toward the proteasome. Abnormal activation of NF-kB followed by the stimulation of cytokine synthesis has been observed in a variety of inflammatory and infectious diseases. Activation of NF-kB is also essential for angiogenesis and for expression of adhesion molecules (CAMs and selects), thus proteasome inhibitors may also have utility in the treatment of diseases associated with the vascular system.

It is well documented that the ubiquitin-proteasome pathway is critical for the regulated destruction of cyclins that govern the exit from mitosis and allow cells to progress into the next phase of the cell cycle (Glotzer, M., Murray, A.W. and Kirschner, M.W. (1991)

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Nature 349, 132-138). Thus, inhibiting the degradation of cyclins by using proteasome inhibitors causes growth arrest. Therefore another potential utility of proteasome inhibitors is their use in the treatment of diseases that result from abberrant cell division.

Several classes of peptidic inhibitors of 20S proteasome have been reported in the recent literature. The α-ketoamide group has been used in protease inhibitors for numerous indications. Specifically, a series of α-ketocarbonyl and boronic ester derived dipeptides (Iqbal, M., Chatterjee, S., Kauer, J.C., Mallamo, J.P., Messina, P.A., Reiboldt, A. and Siman, R. 1996 *Bioorg. Med.Chem. Lett* 6, 287-290) have been reported as potent inhibitors of 20S proteasomal function. Derivatives of 3-indolepyruvic acid have been claimed as pharmaceutically active compounds for the treatment of disturbances of the central nervous system (De Luca, et al WO 88/09789) through a mechanism that modulates kynurenic acid levels in the brain.

Even though various compositions have been discovered that inhibit cell proliferation to some degree, there remains a need for more potent compounds that inhibit cell proliferation via the 20S proteasome.

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SUMMARY OF THE INVENTION

It is an object of this invention to provide a method for inhibiting cell proliferation in mammals that uses a therapeutically effective amount of a composition heretofore unknown for its cell proliferative inhibition properties.

It is also an object of this invention to provide a method for the treatment of diseases that may be affected by the inhibition of proteosomal function.

Further, it is an object of this invention to provide a method for the treatment of proliferative diseases that operates by inhibiting proteasomal function.

It is another object of this invention to use a therapeutically effective amount of the compositions described herein to inhibit cell proliferative disorders in humans.

Yet another object of this invention is the use of a therapeutically effective amount of the compositions described herein to inhibit proteasomal function.

In one embodiment, this invention is a composition of matter having the formula:

$$X_2$$
 \mathbb{N}
 \mathbb{N}
 \mathbb{N}
 \mathbb{N}
 \mathbb{N}

wherein X₂ is Ar or Ar-X₃ wherein X₃ is -C=O, or -CH₂CO-, and wherein Ar is phenyl, substituted phenyl, indole, substituted indoles, and any other heteroaryls; R₁, and R₂ are each individually selected from the side chains of the known natural α-amino acids and unnatural amino acids, hydrogen, 1-10 carbon linear and branched alkyl, 1-10 carbon linear and branched substituted alkyl, aryl, substituted aryl, 1-10 carbon linear, branched substituted aryl, alkoxyaryl, 3-8 carbon cycloalkyl, heterocycle substituted heterocycle, heteroaryl and

substituted heteroaryl; X₁ is selected from hydroxide, monoalkylamino, dialkylamino, alkoxide, arylkoxide and

wherein X_4 is hydroxide, arylamino, monoalkylamino, dialkylamino, alkoxide, or arylalkoxide; and

 R_3 is selected from the known natural α -amino acids, unnatural amino acids, hydrogen, 1-10 carbon linear and branched alkyl, 1-10 carbon linear and branched substituted alkyl, aryl, substituted aryl, 1-10 carbon linear and branched substituted aryl, alkoxyaryl, 3-8 carbon cycloalkyl, heterocycle, substituted heterocycle, heteroaryl and substituted heteroaryl.

In another embodiment, this invention is a method for inhibiting proteasomal protease factor in mammals comprising administering a therapeutically effective amount of the composition disclosed above to the mammal.

In still another embodiment, this invention is a pharmaceutical composition of matter comprising the composition of claim 1 and one or more pharmaceutical excipients.

DESCRIPTION OF THE CURRENT EMBODIMENT

The invention is a method for inhibiting cell proliferation disorders, infectious diseases, and immunological diseases in mammals, and especially in humans, using compositions having the following general formula:

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$$X_2$$
 R_1
 R_2
 R_1

where:

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 X_2 is Ar or Ar- X_3 wherein X_3 is -C=O, -CH₂CO-, or (CH₂)n where n=0-2 and wherein Ar is phenyl, substituted phenyl, indole, substituted indoles, and any other heteroaryl.

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 R_1 , and R_2 are each individually selected from the side chains of the known natural α -amino acids and unnatural amino acids: hydrogen, 1-10 carbon linear and branched alkyl, 1-10 carbon linear and branched substituted alkyl, aryl and substituted aryl, 1-10 carbon linear and branched substituted aryl, alkoxyaryl, 3-8 carbon cycloalkyl, heterocycle and substituted heterocycle, or heteroaryl and substituted heteroaryl. R_2 is preferably biaryl or biphenyl. R_1 is preferably isobutyl. R_2 is selected from -OH, mono or dialkylamino, alkoxide, arylkoxide and

wherein:

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X₄ is -OH, arylamino, mono or dialkylamino, alkoxide, or arylalkoxide; and preferably -OH

10 R_3 is selected from the side chains of known natural α -amino acids and unnatural amino acids, hydrogen, 1-10 carbon linear alkyl and branched alkyl substituents,

1-10 carbon linear and branched substituted alkyl, aryl and substituted aryl, 1-

carbon linear and branched substituted aryl, alkoxyaryl, 3-8 carbon cycloalkyl, heterocycle and substituted heterocycle, or heteroaryl and substituted heteroaryl.

R3 is preferably CO₂H, CH₂CO₂H, (CH₂)₂CO₂H, Arg, Lys, Asn, Gln, Asp, Glu, Phe, and Nle.

The following are definitions for certain terms used herein.

"Halogen" refers to fluorine, bromine, chlorine, and iodine atoms.

"Hydroxyl" refers to the group -OH.

"Thiol" or "mercapto" refers to the group -SH.

"alkyl" refers to a cyclic, branched or straight chain, alkyl group of one to ten carbon atoms. This term is further exemplified by such groups as methyl, ethyl, n-propyl, i-propyl, n-butyl, t-butyl, i-butyl (or 3-methylpropyl), cyclopropylmethyl, i-amyl, n-amyl, n-hexyl and

5 the like.

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"Substituted alkyl" refers to lower alkyl as just described including one or more groups such as hydroxyl, thiol, alkylthiol, halogen, alkoxy, amino, amido, carboxyl, cycloalkyl, substituted cycloalkyl, heterocycle, cycloheteroalkyl, substituted cycloheteroalkyl, acyl, carboxyl, aryl, substituted aryl, aryloxy, hetaryl, substituted hetaryl, aralkyl, heteroaralkyl, alkyl alkenyl, alkyl alkynyl, alkyl cycloalkyl, alkyl cycloheteroalkyl, cyano. These groups may be attached to any carbon atom of the lower alkyl moiety.

"Aryloxy" denotes groups -OAr, where Ar is an aryl, substituted aryl, heteroaryl, or substituted heteroaryl group as defined below.

"Amino" denotes the group NRR', where R and R' may independently be hydrogen, lower alkyl, substituted lower alkyl, aryl, substituted aryl, hetaryl, or substituted hetaryl as defined below or acyl.

"Amido" denotes the group -C(O)NRR', where R and R' may independently by hydrogen, lower alkyl, substituted lower alkyl, aryl, substituted aryl, hetaryl, substituted hetaryl as defined below.

"Carboxyl" denotes the group -C(O)OR, where R may independently be hydrogen, lower alkyl, substituted lower alkyl, aryl, substituted aryl, hetaryl, substituted hetaryl and the like as defined.

"Aryl" or "Ar" refers to an aromatic carbocyclic group having at least one aromatic ring (e.g., phenyl or biphenyl) or multiple condensed rings in which at least one ring is aromatic, (e.g., 1,2,3,4-tetrahydronaphthyl, naphthyl, anthryl, or phenanthryl).

"Substituted aryl" refers to aryl optionally substituted with one or more functional groups, e.g., halogen, lower alkyl, lower alkoxy, alkylthio, acetylene, amino, amido, carboxyl, hydroxyl, aryl, aryloxy, heterocycle, hetaryl, substituted hetaryl, nitro, cyano, thiol, sulfamido and the like.

"Heterocycle" refers to a saturated, unsaturated, or aromatic carbocyclic group having a single ring (e.g., morpholino, pyridyl or furyl) or multiple condensed rings (e.g., naphthpyridyl, quinoxalyl, quinolinyl, indolizinyl or benzo[b]thienyl) and having at least one hetero atom, such as N, O or S, within the ring, which can optionally be unsubstituted or substituted with, e.g., halogen, lower alkyl, lower alkoxy, alkylthio, acetylene, amino, amido, carboxyl, hydroxyl, aryl, aryloxy, heterocycle, hetaryl, substituted hetaryl, nitro, cyano, thiol, sulfamido and the like.

"Heteroaryl" or "hetar" refers to a heterocycle in which at least one heterocyclic ring is aromatic. Preferred heteroaryls are phenyl, substituted phenyl, indole and substituted indoles.

"Substituted heteroaryl" refers to a heterocycle optionally mono or poly substituted with one or more functional groups, e.g., halogen, lower alkyl, lower alkoxy, alkylthio, acetylene, amino, amido, carboxyl, hydroxyl, aryl, aryloxy, heterocycle, hetaryl, substituted hetaryl, nitro, cyano, thiol, sulfamido and the like.

"Cycloalkyl" refers to a divalent cyclic or polycyclic alkyl group containing 3 to 15 carbon atoms.

"Substituted cycloalkyl" refers to a cycloalkyl group comprising one or more substituents with, e.g., halogen, lower alkyl, substituted lower alkyl, alkoxy, alkylthio, acetylene, aryl, aryloxy, heterocycle, hetaryl, substituted hetaryl, nitro, cyano, thiol, sulfamido and the like.

Examples of compounds that may be useful in the therapeutic methods of this invention, and specifically, useful as inhibitors of proteosomal function, are identified in Table 1 below:

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Examples of compounds that may be useful in the therapeutic method of this invention (specifically, useful as inhibitors of proteosomal function) are listed in the table below:

Tabl	le I. Com	position	s used to inhibit 20	S proteosor	ne		
*	Ar	X2	R ₂	R ₁	\mathbf{x}_1	R ₃	X ₃
1	phenyl	сн₂со	**	T)	ОН		
2	Indole	сн₂со	~~~		ОН		
3	Indole	СН₂СО		T	PhCH ₂ N		
4	Indole	со		T		z-\{	он
5	Indole	сн₂со		Y		}_i_{	он
6	phenyl	СН₂СО				1 -\	он
7	phenyl	сн₂со				\\	PhCH ₂ N
8	Indole	со				}-z-	PhCH ₂ N

9	Indole	СН₂СО	1 1	~	 	PhCH ₂ N
10	Indole	со		ОН		
11	phenyl	со	***	ОН		
12	phenyl	со			\	ОН
13	phenyl	СН2СО				он
14	Indole	со				он
15	Indole	CH ₂ CO	* 	~		ОН
16	phenyl	со				ОН
17	indole	Сн₂со				ОН
18	Indole	СН₂СО				ОН

19	Indole	СН₂СО			ОН
20	Indole	CH₂CO		ÇH₃ ✓	ОН
21	indole	СН₂СО		OH	он
22	Indole	CH ₂ CO	$\left \begin{array}{c} \downarrow \\ \downarrow \end{array}\right $		он
23	Indole	CH ₂ CO			он
24	Indole	СН₂СО			ОН
25	Indole	сн₂со		SH,	он
26	indole	СН2СО		ОН	он
27	Indole	СН₂СО			ОН
28	indole	СН₂СО	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		ОН
29	indole	сн₂со			ОН

30	Indole	СН₂СО		 ~		CH₃	ОН
31	Indole	CH ₂ CO		\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		ОН	он
32	indole	сн₂со		~			ОН
33	Indole	Сн₂со	*			.	он
34	Indole	сн₂со		 			ОН
35	Indole	СН₂СО		 		ÇH ₃	ОН
36	Indole	Сн₂СО		\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		~~	он
37	Indole	СН₂СО	**************************************	7			ОН
38	Indole	CH ₂ CO	β-Ala	$ \gamma $		☆ Y	ОН
39	Indole	СН₂СО		~		. ★	ОН
4 0	Indole	Сн₂со		~		* 	ОН
		1.		1	-	! ~	On

41	Indole	CH ₂ CO	№ ОН	Y	ОН
42	Indole	CH ₂ CO	CH₃		ОН
43	Indole	СН₂СО	HO ₂ C		ОН
44	Indole	CH ₂ CO	\mathcal{X}		ОН
45	Indole	СН ₂ СО		\downarrow	ОН
46	Indole	СН2СО	⋄ Он		он
47	Indole	СН₂СО	.		ОН
48	Indole	СН2СО		$\left \begin{array}{c} \sim \\ \sim \end{array}\right $	Он
49	Indole	CH ₂ CO			ОН
50	Indole	Сн₂со	•		ОН

51	Indole	Сн₂со				он
52	Indole	сн₂со	↓		\\	он
53	Indole	сн₂со	В-А1а		\r_\	он
54	Indole	CH ₂ CO		$ \uparrow $	γ-z	он
55	Indole	СН₂СО			}-z	он
56	Indole	СН₂СО	№ он		\	он
57	Indole	Сн₂со	CH₃		 *	он
58	Indole	СН2СО	HO ² C		\	он
59	Indole	СН₂СО	\mathcal{X}	T	\	он
60	Indole	сн₂со			\\	ОН

61	Ind le	сн₂со	ОН		\	ОН
62	Indole	сн₂со			}-I	он
63	Indole	CH ₂ CO			\ r	он
64	Indole	сн₂со			}-z	он
65	Indole	СН₂СО			}-r	он
66	Indole	сн₂со		T	}-I	он
67	Indole	сн₂со	}-z	T	ÇH₃	он
68	Indole	СН₂СО	β-Ala		ÇH₃ ✓	он
69	Indole	СН ₂ СО			ÇH₃	он
70	Indole	CH ₂ CO		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	ÇH₃	он

71	Indole	СН₂СО	ОН	T	CH₃	Он	1
72	Indole	сн₂со	СH ₃	$\left \gamma \right $	CH₃	ОН	
73	Indole	Сн₂со	HO₂C →		CH₃	ОН	
74	Indole	СН₂СО	\downarrow		CH₃	ОН	
75	Indole	СН₂СО			CH ₃	ОН	
76	indole	Сн ₂ СО	→ OH		CH₃ ✓	ОН	
77	Indole	СН₂СО	•		CH ₃	ОН	
78	Indole	СН₂СО		~	 CH₃	Он	
79	ind ole	Сн2СО	.		CH₃	Он	
80	indole	СН₂СО	•		CH ₃	ОН	!
				1			

81	Indole	сн₂со			5	СH ₃	он
82	Indole	CH ₂ CO	\	T		ОН	он
83	Indole	сн₂со	₩ он			ОН	он
84	Indole	СН₂СО	β-Ala		1.	ОН	он
85	Indole	со			~ ~ ~		ОН
86	Indole	СН₂СО			7	·	ОН
87	Indole	сн₂со				ОН	ОН
88	Indole	сн₂со				ОН	ОН
89	Indole	СН ₂ СО	\-\rm\rm\rm\rm\rm\rm\rm\rm\rm\rm\rm\rm\rm\		HO.	<u>)</u>	ОН
90	Indole	СН₂СО	β-Ala		но	$\sum_{i=1}^{\infty}$	ОН

91	Indole	СН₂СО		T	но)	ОН	
92	Indole	СН₂СО			нс) }	ОН	
93	Indole	СН₂СО	№	T	но	~ ``	ОН	
94	Indole	CH ₂ CO	CH₃		но	<u>)</u>	ОН	
95	Indole	CH ₂ CO	HO ₂ C		но	<u>)</u>	он	
96	indole	СН ₂ СО	\mathcal{X}		НО	\hat{J}	он	
97	Indole	Сн ₂ СО			но),	ОН	
98	Indole	СН₂СО	→ OH		но,)	ОН	
99	Indole	Сн₂со			но		ОН	
100	Indole	СН₂СО			но		ОН	
101	Indole	сн₂со			но		он	
) (1	~~	1	1			

10	Indole	СН₂СО		T	HO	он
10	Indole	Сн₂со			₩O >	ОН
10	4 Indole	СН2СО	\	T		он
10	5 Indole	СН₂СО	β-А1а			он
10	Indole	CH ₂ CO				ОН
10	Indole	СН₂СО		$\left \begin{array}{c} 1 \\ 1 \end{array} \right $		ОН
100	Indole	сн₂со	№ он			ОН
109	indole	СН₂СО	ÇH₃ ✓		.	он
110	Indole	СН₂СО	HO ₂ C			Он
111	indole	СН₂СО	Val			ОН

112	Indol	СН₂СО	Nva		он
113	Indole	снұсо	₩ OH		он
114	Indole	СН₂СО			он
115	indole	СН₂СО			он
116	Indole	СН₂СО			он
117	Indole	сн₂со			он
118	Indole	Сн₂со			ОН
119	Indole	СН₂СО	\r		ОН
120	Indole	Сн₂со	β-Ala	T	ОН

121	Indole	СН₂СО		T_	он
122	Indole	СН ₂ СО			он
123	Indole	СН₂СО	~ oн _		он
124	Indole	СН2СО	СН		ОН
125	Indole	СН₂СО	HO₂C		он
126	indole	СН₂СО	\mathcal{X}		он
127	indole	СН₂СО			он
128	Indole	СН₂СО	₹ OH		ОН
129	Indole	СН₂СО	1	T	ОН

130	Indole	CH ₂ Cc		17		_	Он	i
131	Indole	Сн₂Сс					ОН	
132	indole	CH ₂ CC					Он	
133	Îndole	CH ₂ CC					Он	
134	Indole	СН₂СО		T		HO ₂ C	ОН	
135	Indole	CH ₂ CO	β-Ala			HO ₂ C	ОН	
136	Indole	сн₂со			-	HO ₂ C	ОН	
137	Indole	CH ₂ CO		~		HO ₂ C	ОН	
138	Indole	СН₂СО	С Дон			HO ₂ C	ОН	
139	Indole	СН2СО	CH3 ◆			HO ₂ C	он	
i 	I (l i		1 1	l	• •		

140	Indole	СН₂СО	HO ₂ C	1	HO ₂ C	Он
141	Indole	СН₂СО			HO ₂ C	он
142	Indole	сн₂со			HO ₂ C	он
143	Indole	СН₂СО	ОН	$\left \stackrel{\uparrow}{ \swarrow} \right $	HO ₂ C	он
144	Indole	СН₂СО		$ \gamma $	HO ₂ C	он
145	Indole	CH ₂ CO		$\left \begin{array}{c} \uparrow \\ \uparrow \end{array} \right $	HO ₂ C	он
146	Indole	СН₂СО	· N		HO ₂ C	он
147	Indole	СН ₂ СО			HO ₂ C \searrow	ОН
148	Indole	сн₂со	4.4'-BPA		HO ₂ C	ОН
149	Indole	сн₂со	}-I	$ \gamma^{\dagger} $	СОЙ	ОН
150	Indole	СН₂СО	β-Ala		CO⁵H	ОН

151	Indole	Сн₂со		$ \Upsilon $		СОЗН	он
152	Indole	Сн2СО				ÇO₂H	ОН
153	Indole	Сн₂со	Дон			CO⁵H	Он
154	Indole	СН₂СО	CH ₃			СОРН	Он
155	Indole	СН₂СО	HO₂C →			CO₂H	ОН
156	Indole	CH ₂ CO	\downarrow			СО₂Н	ОН
157	Indole	CH ₂ CO				CO₂H	он
158	Indole	СН₂СО	→ OH			CO2H.	он
159	Indole	Сн₂со	* -			ÇO₂H	он
160	Indole	СН₂СО			·	ÇO₂H	ОН
161	indole	СН₂СО				√ CO2H	ОН
			~~			*	

162	indole	СН₂СО		$ \zeta $	СОЭН	ОН
163	Indole	Сн₂со			ÇO₂H	ОН
164	Indole	сн₂со		$\left \gamma \right $	\downarrow	он
165	indole	Сн₂СО		$ \Upsilon $	\searrow	ОН
167	indole	СН₂СО	\downarrow	$ \chi $	\mathcal{X}	Он
168	indole	CH ₂ CO			$\stackrel{\smile}{\searrow}$	ОН
169	Indole	CH ₂ CO		$ \chi $	\ -I	ОН
170	Indole	СН₂СО			_	ОН
1	į	1	♣			

171	Ind le	сн₂со				Он
172	Indole	СН₂СО				он
173	Indole	СН₂СО	\$ \(\sigma_z \)			он
174	Indole	Сн₂СО				ОН
175	Indole	СН₂СО				Он
176-	Indole	Сн₃со		~	HO ₂ C	ОН
177	Indole	Сн₂СО			CO⁵H	ОН
178	Indole	Сн₂со				ОН

179	Indole	сн₂со					ОН
180	Indole	CH ₂ CO				HO₂C .	ОН
181	Indole	СН₂СО		T	·	СОЭН	ОН
182	Indole	СН₂СО	*	*		он	
183	Indole	СН₂СО	}-x	H		\	ОН
184	Indole	СН₂СО	}-I	$\left \overset{\bullet}{\widetilde{\chi}} \right $		\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	он
185	Indole	Сн₂СО				H ₂ NOC →	ОН
186	Indole	сн₂со		\downarrow		CONH₂	ОН
-	•		i		j		1

187	Indole	сн₂со		T			ОН
188	Indole	СН₂СО				β-Alanine	ОН
189	Indole	СН₂СО		~			Он
190	Indole ,	СН₂СО			·		ОН
191	Indole	CH ₂ CO			·	HN Y NH2	- ОН
192	Indole	сн₂со			-		он
193	Indole	СН2СО		$ \tau $		NH ₂	он
	Ì		~			~	

194	Indole	сн₂со				ОН
195	Indole	Сн ₂ СО			HO ₂ C	он
196	Indole	СН₂СО		~	₩ 1	он
197	Indole	СН₂СО			HN → NH2	ОН
198	Indole	сн₂со			s N	ОН
199	Indole	СН2СО		*	$\stackrel{\sim}{\downarrow}$	он
200	Indole	сн₂со	HO ₂ C		*	он
201	Indole	СН₂СО		\downarrow	*	ОН

202	Indole	СН₂СО	ОН	$ \mathcal{X} $	$ \chi$	Он
203	indole	сн₂со		~		он
204	indole	сн₂со		$\widehat{\mathcal{T}}$	N T T T T T T T T T T T T T T T T T T T	ОН
205	Indole	сн₂со		T_	HN YNH2	он
206	indole	сн₂со				он -
207	Indole	сн₂со	B-Alanine	β-Alanine	ОН	

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The compounds described above are useful for treating diseases and disorders mediated by the 20S proteasome such as antiproliferative diseases, cancer, inflammation. It is preferred that the compositions of this invention are used to treat antiproliferative disorders and inflammation. It is most preferred that the compounds of this invention are used to treat inflammatory diseases.

The compounds of the present invention are useful for treating disorders mediated by 20S proteasome in mammals.

The compounds of this invention may be administered to mammals both prophylactically and therapeutically by any administration protocol that is capable of supplying at least one compound of this invention to a 20S proteasome. Non-limiting examples of useful administration protocols include orally, parenterally, dermally, transdermally, rectally, nasally or by any other suitable pharmaceutical composition administration protocol that is within the knowledge of one skilled in the art.

The compositions of this invention may be administered in suitable pharmaceutical dosage forms. The pharmaceutical dosage form will depend largely upon the administration protocol used. The term pharmaceutical dosage form refers to items such as tablets, capsules, liquids and powders, comprising 20S proteasome inhibitors of this invention alone or in the presence of one or more pharmaceutical excipients. The choice of additives such as excipients and adjuvants again will depend largely upon the chosen administration protocol. Those skilled in the pharmaceutical arts will recognize a wide variety of formulations and vehicles for administering compositions of this invention.

The administration protocol chosen for compounds of this invention will ultimately dictate the final form and composition of pharmaceutical dosage forms comprising the 20S proteasome inhibitors of this invention. For example, internal administration of compounds of this invention is effected, orally, in the form of powders, tablets, capsules, pastes, drinks,

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granules, or solutions, suspensions and emulsions which can be administered orally, or bolus, in medicated food, or in drinking water. Internal administration may also be accomplished using a timed release formulation including additives such as surfactant or starch coated capsules, or using a quick release formulation such as a freeze-dried fast dissolving tablet. Dermal administration is effected, for example, in the form of transdermal patches, spraying or pouring-on and spotting-on. Parenteral administration is effected, for example, in the form of injection (intramuscularly, subcutaneously, intravenously, intraperitoneally) or by implants.

Suitable pharmaceutical dosage forms incorporating the 20S proteasome inhibitors of this invention include but are not limited to solutions such as solutions for injection, oral solutions, concentrates for oral administration after dilution, solutions for use on the skin or in body cavities, pour-on and spot-on formulations, gels; emulsions and suspension for oral or dermal administration and for injection; semi-solid preparations; formulations in which the active compound is incorporated in cream base or in an oil-in-water or water-in-oil emulsion base; solid-preparations such as powders, premixes or concentrates, granules, pellets, tablets, boli, capsules; aerosols and inhalants, and shaped articles containing active compound.

Pharmaceutical dosage forms that are solutions may be administered by injection intravenously, intramuscularly and subcutaneously. Solutions for injection are prepared by dissolving the active compound in a suitable solvent and, if appropriate, adding adjuvants such as solubilizers, acids, bases, buffer salts, antioxidants and preservatives. The solutions are sterile-filtered and drawn off.

Alternatively, solutions including compositions of this invention may be administered orally. Concentrates of compositions of this invention are preferably administered orally only after diluting the concentrate to the administration concentration. Oral solutions and concentrates are prepared as described above in the case of the solutions for injection. Solutions for use on the skin are applied dropwise, brushed on, rubbed in, splashed on or

sprayed on. These solutions are prepared as described above in the case of solutions for injection.

Gels are applied to the skin, or introduced into body cavities. Gels are prepared by treating solutions which have been prepared as described in the case of the solutions for injection with such an amount of thickener that a clear substance of cream-like consistency is formed, or by any other means known to one skilled in the art.

Pour-on and spot-on formulations are poured onto, or splashed onto, limited areas of the skin, the active compound penetrating the skin and acting systemically. Pour-on and spot-on formulations are prepared by dissolving, suspending or emulsifying the active compound in suitable solvents or solvent mixtures which are tolerated by the skin. If appropriate, other adjuvants such as colorants, resorption accelerators, antioxidants, light stabilizers, and tackifiers are added.

Emulsions can be administered orally, dermally or in the form of injections. Emulsions are either of the water-in-oil type or of the oil-in-water type. They are prepared by dissolving 20S proteasome inhibitors either in the hydrophobic or in the hydrophilic phase and homogenizing the phase with a solvent of the opposite phase with the aid of suitable adjuvants such as emulsifiers, colorants, resorption accelerators, preservatives, antioxidants, light stabilizers, and viscosity-increasing substances.

Suspensions can be administered orally, dermally or in the form of injection. They are prepared by suspending the active compound in a liquid if appropriate with the addition of further adjuvants such as wetting agents, colorants, resorption accelerators, preservatives, antioxidants and light stabilizers.

The pharmaceutical compositions of this invention may include one or more additives in the form of pharmaceutically acceptable additives. Useful additives include solvents, solubilizers, preservatives, thickeners, wetting agents, colorants, resorption accelerators,

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antioxidants, light stabilizers, tackifiers, viscosity increasing substances, fillers, flavorings, lubricating agents, and any other pharmaceutical composition additive known to those skilled in the art.

The additive may be a solvent such as water, alcohols such as ethanol, butanol, benzyl alcohol, glycerol, propylene glycol, polyethylene glycols, N-methyl-pyrrolidone, alkanols, glycerol, aromatic alcohols such as benzyl alcohol, phenylethanol, phenoxyethanol, esters such as ethyl acetate, butyl acetate, benzyl benzoate, ethers such as alkylene glycol alkyl ethers such as dipropylene glycol mono-methyl ether, diethylene glycol mono-butyl ether, ketones such as acetone, methyl ethyl ketone, aromatic and/or aliphatic hydrocarbons, vegetable or synthetic oils, DMF, dimethylacetamide, N-methyl-pyrrolidone, 2,2-dimethyl-4-oxy-methylene-1,3-dioxolane.

The following additives may be useful as solubilizers of the compositions of this invention: solvents which enhance solution of the active compound in the main solvent or which prevent its precipitation. Examples are polyvinylpyrrolidone, polyoxyethylated castor oil, polyoxyethylated sorbitan esters.

Useful preservatives are, for example, benzyl alcohol, trichlorobutanol, p-hydroxybenzoic esters, and n-butanol.

Useful thickeners include inorganic thickeners such as bentonite, colloidal silica, aluminum monostearate, organic thickeners such as cellulose derivatives, polyvinyl alcohols and their copolymers, acrylates and methacrylates.

Other liquids which may be useful in pharmaceutical dosage forms of this invention are, for example, homogeneous solvents, solvent mixtures, and wetting agents which are typically surfactants.

Useful colorants are all colorants which are non-toxic and which can be dissolved or suspended.

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Useful resorption accelerators are DMSO, spreading oils such as isopropyl myristate, cipropylene glycol pelargonate, silicone oils, fatty acid esters, triglycerides, fatty alcohols.

Useful antioxidants are sulphites or metabisulphites such as potassium metabisulphite, ascorbic acid, butylhydroxytoluene, butylhydroxyanisole, tocopherol.

A useful light stabilizer is novantisolic acid.

Useful tackifiers include cellulose derivatives, starch derivatives, polyacrylates, natural polymers such as alginates, gelatin.

Useful emulsifiers include non-ionic surfactants such as polyoxyethylated castor oil, polyoxyethylated sorbitan monooleate, sorbitan monostearate, glycerol monostearate, polyoxyethyl stearate, alkylphenol polyglycol ethers; ampholytic surfactants such as Di-Na N-lauryl- beta -iminodipropionate or lecithin; anionic surfactants, such as Na-lauryl sulphate, fatty alcohol ether sulphates, the monoethanolamine salt of mono/dialkylpolyglycol ether orthophosphoric esters; cationic surfactants such as cetyltrimethylammonium chloride.

Useful viscosity-increasing substances and substances which stabilize a therapeutic emulsion include carboxymethylcellulose, methylcellulose and other cellulose and starch derivatives, polyacrylates, alginates, gelatin, gum Arabic, polyvinylpyrrolidone, polyvinyl alcohol, copolymers of methyl vinyl ether and maleic anhydride, polyethylene glycols, waxes, colloidal silica or mixtures of the substances mentioned.

To prepare solid pharmaceutical dosage forms, the active compound is mixed with suitable additives, if appropriate with the addition of adjuvants, and the mixture is formulated as desired. Examples of physiologically acceptable solid inert additives include sodium chloride, carbonates such as calcium carbonate, hydrogen carbonates, aluminum oxides, silicas, clays, precipitated or colloidal silicon dioxide, and phosphates. Examples of solid organic additives include sugars, cellulose, foods such as dried milk, animal meals, cereal meals and coarse cereal meals and starches. Other suitable additives include lubricants and

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gliding agents such as magnesium stearate, stearic acid, talc, bentonites; disintegrants such as starch or crosslinked polyvinylpyrrolidone; binders such as, starch, gelatin or linear polyvinylpyrrolidone; and dry binders such as microcrystalline cellulose.

In the pharmaceutical dosage forms described herein, the active compounds can be present in the form of a mixture with at least one other 20S proteasome inhibitor. Alternatively, or in addition, the pharmaceutical dosage forms of the invention can, in addition to at least one 20S proteasome inhibitor, include any pharmaceutical compound that is capable of treating any known malady or disorder where the administration of both together create no unacceptable adverse effects.

Methods for treating 20S proteasome mediated diseases and disorders comprises the administration of an effective quantity of the chosen compound or combinations thereof, preferably dispersed in a pharmaceutical dosage form. Ready-to-use pharmaceutical dosage forms of this invention contain the active compound in concentrations of from 10 ppm to 20 per cent by weight, and preferably of from 0.1 to 10 per cent by weight. Pharmaceutical dosage forms of this invention that are diluted prior to administration, preferably contain the active compound in concentrations of from 0.5 to 90 per cent by weight, and preferably of from 5 to 50 per cent by weight. In general, it has proved advantageous to administer amounts of approximately 0.01mg to approximately 100 mg of active compound per kg of body weight per day to achieve effective results.

The amount and frequency of administration of pharmaceutical dosage forms comprising 20S proteasome inhibitors of this invention will be readily determined by one skilled in the art depending upon, among other factors, the route of administration, age and condition of the patient. These dosage units may be administered one to ten times daily for acute or chronic disease. No unacceptable toxicological effects are expected when compounds of the invention are administered in accordance with the present invention.

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The pharmaceutical dosage forms comprising 20S proteasome inhibitors of this invention are made following the conventional techniques of pharmacy involving milling, mixing, granulation, and compressing, when necessary, for tablet forms; or milling, mixing and filling for hard gelatin capsule forms. When a liquid additive is used, the preparation will be in the form of a syrup, elixir, emulsion or an aqueous or non-aqueous suspension. Such a liquid formulation may be administered directly p.o. or filled into a soft gelatin capsule.

While the compositions described herein may be administered as described above, it is preferred that the method of this invention is achieved by administering the compound described herein orally. When the oral administration route is chosen, a larger quantity of reactive agent will be required to produce the same effect as a smaller quantity given for example parenterally. In accordance with good clinical practice, it is preferred to administer the compound according to this method at a concentration level that would produce effective therapeutic results without causing any harmful side effects.

The compositions of this invention have non-therapeutic utility as well. The compositions of this invention are useful as analytical standards for 20S proteasome inhibitor assays.

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Example 1

The compounds useful in the therapeutic method of this invention are prepared by conventional methods of organic chemistry. References that may be consulted in describing the art of the synthesis of these compounds include Bodansky's "The Practice of Peptide Synthesis," Springer-Verlag, First Edition, 1984; "Protective Groups in Organic Synthesis," Second Edition, John Wiley and Sons, New York, 1991. All peptide couplings are accomplished at room temperature with gentle and constant agitation. Peptide couplings and deprotections are monitored using the Kaiser test for amines. Xaa refers to any of the commercially available amino acids that may be purchased pre-attached to the MBHA resin. Yaa and Zaa refer to any of the commercially available amino acids.

The compounds of this invention may be prepared by solid phase peptide synthesis (SPPS) in the general procedure that follows: Xaa-MBHA-resin is weighed and transferred to a syringe equipped with a fritted filter. The resin is pre-swollen in DMF and then the N-terminal protecting group is removed by treatment with 30%, piperidine in DMF for 30 minutes. The deprotection solution is removed. The deprotected resin is washed five times with DMF, five times with MeOH, and then five times with DMF. Amino acid Yaa may then be coupled to the deprotected resin using a solution of Yaa in DMF containing 3 equivalents each of Yaa, carbodiimide coupling reagent and HOBT (hydroxy benzotriazole). Succesive couplings with solutions of Yaa may be necessary to achieve coupling efficiencies that pass the Kaiser test. The N-terminal group deprotection and Yaa coupling step may be repeated to couple a third amino acid Zaa. The final coupling step uses ketoacid, carbodiimide, and HOBT in DMF, and this step is repeated until the coupling passes the Kaiser test. The completed peptide sequence on resin is dried under vaccuum for at least six hours and then cleaved by treatment for 2.5 hours with either 95/5 trifluoroacetic acid/water or a freshly prepared solution of 90%, trifluoroacetic acid, 3% ethanedithiol, 5% thioanisole, and 2%

anisole. The cleaved products are recovered by either lyophillization from water or trituration from diethyl ether. Product purities are estimated from TLC. Selected peptide samples are checked by ¹H NMR to confirm product identity.

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Example 2

In this Example (3'-Indolepyruvic acid)- N-biphenylalanine-D-Leu-Asp-OH was prepared according to the method of Example 1.

Fmoc-N-Asp(Ot-Bu)-MBHA-resin (20 mg) is weighed and transferred to a syringe equipped with a fritted filter. The resin is pre-swollen in 1 mL DMF for 30 minutes. The Fmoc (fluorenylmethyloxycarbonyl) protecting group is removed by treatment with 20% piperidine in DMF for 30 minutes. The deprotection solution is removed. The deprotected resin is washed five times with DMF, five times with MeOH, and then five times with DMF. Fmoc-D-Leu-OH is coupled to the deprotected resin (1eq) using a solution of Fmoc-D-Leu-OH (3 eq) in 1 mL DMF containing carbodiimide (3 eq) and HOBT (hydroxy benzotriazole) (3 eq). A second or third coupling with solutions of Fmoc-D-Leu-OH may be necessary to achieve coupling efficiencies that pass the Kaiser test. The Fmoc deprotection and amino acid coupling step are repeated to couple Fmoc-N- (4,4-biphenyl)alanine. The final coupling step uses indolepyruvic acid (5eq), diisopropylcarbodiimide (5eq), and HOBT (5eq) in DMF, and this step is repeated until the coupling passes the Kaiser test. The completed peptide sequence on resin is dried under vacuum for at least six hours and then cleaved by treatment for 2.5 hours with 1 mL of either 95/5 trifluoroacetic acid/water or a freshly prepared solution of 90% trifluoroacetic acid, 5% thioanisole, 3% ethanedithiol, and 2% anisole. The cleaved products are recovered by either lyophillization from water or trituration from diethyl ether. Product purities are estimated from TLC.

¹H NMR (400 MHz, d_6 -DMSO): δ 6.5-7.7 (m, 14H), 4.5 (m, 1H), 4.1(m, 2H), 3.4(m, 2H), 3 (m, H), 2.7 (m, 1H), 1.1-1.5 (m, 3H), 0.5-0.9 (m, 6H).

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Example 3

In this example, (3'-Indolepyruvic acid)-N-biphenylalaanine-D-Leu-Asp-OH was prepared using Chiron Mimotopes Pin Technology

The first amino acid residue Xaa is attached to 4-(hydroxymethyl)phenoxyacetamido handle) resin pins (5.7 µmole/pin) by coupling each pin in 800 µL of coupling solution (100 mM amino acid, 100 mM DIC, 10 mM DMAP, 1/4 DMF/CH₂CI₂) for two hours. The pins are then rinsed with a 5 min DMF wash, two 5 min MeOH washes, and 15 minutes of air drying. Deprotection of the Fmoc group is carried out for 30 min with 800 µL 20% piperidine in DMF. Repeat pin washings (1 DMF wash, 2 MEOH washes, 15 minutes air drying). The second amino acid residue Yaa was coupled (100 mM Yaa, 100 mM DIC, 100 mM HOBT. and bromophenol blue indicator in DMF) until the blue color no longer adheres to the pin surface. The coupling was repeated as necessary. The rinse cycle and Fmoc deprotection washes were then repeated as well. The next amino acid, Zaa, was coupled by repeating the coupling and washing procedures for coupling Yaa, repeating the coupling if necessary. The last residue, indolepyruvic acid is coupled with 15 eq, 100 mM, 15 eq DIC, 15 eq HOBT, and bromophenol blue indicator in DMF. Repeat coupling if necessary. After the last wash, the orange pins were removed from their supports and cleaved in individual 2 mL plastic centrifuge tubes with 1.5 mL of a freshly prepared solution of 90%, trifluoroacetic acid, 5% thioanisole, 3% ethanedithiol, and 2% anisole for 2.5 hours. The pins were removed from the tubes and the mixture was blown to near dryness under a nitrogen stream. Triturate with Et,O and spin down each tube. This step was repeated three times per tube. The precipitated peptides were collected, lyophillized, weighed, and used. Product purity was estimated by TLC. Initial products were cospotted and checked against authentic samples obtained in Example 1.

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Example 4

Compounds of this invention prepared according to the method of Example 1 were tested as follows. The 20S catalytic subunit of the proteasome (also known as the multicatalytic proteinase complex) was purified to homogeneity from bovine brain according to published methods (Wilk S. and Orlowski, M. 1983, 40 842 J.Neurochem). The chymotryptic activity of the complex is measured by the increase in fluorescence following cleavage of the substrate peptide succinyl-leucine-leucine-valine-tyrosine-7-amino-4methyl cournarin. The standard *in vitro* assay consists of 2µg 20S proteasome, 0.1-100µg/ml proteasome inhibitor in 200µl 50mM HEPES, containing 0.1% sodium dodecyl sulphate, pH7.5. The proteolytic reaction is initiated by the addition of 50µM flourogenic peptide substrate and allowed to progress for 15 minutes at 37°C. The reaction is terminated by the addition of 100 µL of 100 mM acetate buffer, pH4.0. The rate of proteolysis is directly proportional to the amount of liberated aminomethylcournarin—which is measured by-fluorescent spectroscopy (EX 370nm, EM 430nm).

The results of the 20S proteasome inhibitor assays are presented in Table II.

Table II.

IC 50 values for the inhibition of the chymotrypsin-like activity of 20S proteasome.

Compound #	IC _{so} μg/mL	Compound #	IC ₅₀ μg/mL
1	10	105	>10
2	10	106	>10
3	>10	107	>10
4	10	108	>10
5	>10	109	>10
6	>10	110	>10
7	>10	111	>10
8	>10	113	>10
9	>10	114	10
10	>10	115	10
11	>10	116	10

Compound #	IC ₅₀ μg/mL	Compound #	IC _{so} μg/mL
12	>10	117	10
13	>10	118	10
14	>10	119	>10
15	10	120	>10
16	10	121	>10
17	>10	122	>10
18	>10	123	>10
19	>10	124	>10
20	>10	125	>10
21	>10	126	>10
22	>10	127	>10
23	>10	128	>10
24	>10	129	10
25	>10	130	10
26	>10	131	10
27	>10	132	
28	>10	133	10
29	>10	134	10
30	>10	135	>10
31	>10	136	>10
32	>10	137	>10
33	>10	138	>10
34	>10	139	>10
35	>10	140	>10
36	>10	141	>10
37	>10	141	>10
38	>10	142	>10
39	>10	143	>10
40	>10	145	10
40	>10	145	10
41	>10		10
43		147	10
43	>10	148	10
45	>10	149	10
	>10	150	>10
46	>10	151	>10
47	>10	152	>10
48	>10	153	>10
49	>10	154	>10
50	>10	155	>10
51	>10	156	>10
52	>10	157	>10
53	>10	158	>10
54	>10	159	>10
55	>10	160	>10

Compound #	IC _{so} μg/mL	Compound #	IC _{sc} μg/mL
56	>10	161	>10 >10
57	>10	162	>10
58	>10	163	
59	>10	164	>10
60	>10	165	>10
61	>10	166	>10
62	>10		>10
63	>10	167	>10
64	>10	168	>10
65	>10	169	>10
66		170	>10
67	>10	171	>10
68	>10	172	>10
	>10	173	>10
69	>10	174	5
70	>10	175	>10
71	>10	176	1
72	>10	177	10
73	>10	178	>10
74	>10	179	>10
75	>10	180	5
76	>10	181	10
77	>10	182	>10
78	>10	183	
79	>10	184	10
80		185	>10
81	>10	186	5
82	>10	187	>10
83	>10	188	>10
84	>10	189	5
85	>10	190	>10
86	>10	191	3 3
87	>10	191	. 3
88	>10	1	3
89		193	>10
90	>10	194	>10
91	>10	195	>10
92	>10	196	>10
92 93	>10	197	10
	>10	198	>10
94 05	>10	199	>10
95 06	>10	200	>10
96 07	>10	201	>10
97	>10	202	>10
98	>10	203	>10
99	>10	204	>10

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Compound #	IC _{so} μg/mL	Compound #	IC ₅₀ μg/mL
100	>10	205	>10
101	>10	206	>10
103	>10	207	10
104	>10		10

Compounds of this invention prepared according to the method of Example 1 were also tested as follows. The 20S catalytic subunit of the proteasome (also known as the multicatalytic proteinase complex) was purified to homogeneity from bovine brain according to published methods (Wilk S. and Orlowski, M 1983, 40 842 J. Neurochem). The tryptic activity of the complex is measured by the increase in fluorescence following cleavage of the substrate peptide CBZ-D-Ala-Leu-Arg-(7-amino -4-methyl cournarin). The standard *in vitro* assay consists of 2µg 20S proteasome, 0.1-100µg/ml proteasome inhibitor in 200ml 50mM HEPES, containing 0.1% sodium dodecyl sulphate, pH 7.5. The proteolytic reaction is initiated by addition of 50mM flurogenic peptide substrate and allowed to progress for 15 minutes at 37°C. The reaction is terminated by the addition of 100 mL of 100 mM acetate buffer, pH4.0. The rate of proteolysis is directly proportional to the amount of liberated aminomethylcournarin which is measured by fluorescent spectroscopy (EX 370nm, EM 430nm). Compounds 1-207 were tested for tryptic activity inhibition and active as inhibitors at >10 µg/mL.

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Example 5

Compounds of this invention prepared according to the method of Example 1 were also tested as follows. The 20S catalytic subunit of the proteasome (also known as the multicatalytic proteinase complex) was purified to homogeneity from bovine brain according to published methods (Wilk S. and Orlowski, M. 1983, 40 842 J. Neurochem). The tryptic activity of the complex is measured by the increase in fluorescene following cleavage of the substrate peptide CBZ D-Ala-Leu-Arg-(7-amino-4-methyl coumarin). The standard *in vitro assay* consists of 20 µg 20S proteasome, 0.1-100µg/ml proteasome inhibitor in 200 µL 50mM HEPES, containing 0.1% sodium dodecyl sulphate, pH 7.5. The proteolytic reaction is initiated by the addition of 50mM fluorogenic peptide substrate and allowed to progress for 15 minutes at 37°C. The reaction is terminated by the addition of 100 µL of 100 mM acetate buffer, pH4.0. The rate of proteolysis is directly proportional to the amount of liberated aminomethylcoumarin which is measured by fluorsecent spectroscopy (EX 370nm, EM 430nm). Compounds 1-207 were tested for tryptic activity inhibition and were active as inhibitors at > 10 µg/mL.

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Example 6

Compounds of this invention prepared according to the method of Example 1 were a so tested as follows. The 20S catalytic subunit of the proteasome (also known as the multicatalytic proteinase complex) was purified to homogeneity from bovine brain according to published methods (Wilk S. and Orlowski, M. 1983, 40 842 J. Neurochem). The tryptic activity of the complex is measured by the increase in fluorescence following cleavage of the substrate peptide CBZ D-Ala-Leu-Glu-(7-amino-4-methyl coumarin). The standard *in vitro assay* consists of 2 µg 20S proteasome, 0.1-100µg/ml proteasome inhibitor in 200ml 50mM HEPES, containing 0.1% sodium dodecyl sulphate, pH 7.5. The proteolytic reaction is initiated by the addition of 50mM fluorogenic peptide substrate and allowed to progress for 15 minutes at 37°C. The reaction is terminated by the addition of mL of 100 mM acetate buffer, pH 4.0. The rate of proteolysis is directly proportional to the amount of liberated aminomethylcoumarin which is measured by fluorescent spectroscopy (EX 370nm, EM 430nm). Compounds 1-207 were tested for peptidylglutamyl activity inhibition at > 10 µg/mL. Compound 190 was active at 5 µg/mL.

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What we claim is:

1. A compound having the formula:

$$X_2$$
 \mathbb{N}
 \mathbb{R}_1
 \mathbb{R}_1

wherein:

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 X_2 is Ar or Ar- X_3 , wherein X_3 is -C=O, -CH₂CO-, or (CH₂)_n where n=0-2, and wherein Ar is phenyl, substituted phenyl, indole, substituted indoles, and any other heteroaryls;

R₁, and R₂ are each individually selected from side chains of the known natural α-amino acids and unnatural amino acids, hydrogen, 1-10 carbon linear and branched alkyl, 1-10 carbon linear and branched substituted alkyl, aryl, substituted aryl, 1-10 carbon linear, branched substituted aryl, alkoxyaryl, 3-8 carbon cycloalkyl, heterocycle substituted heterocycle, heteroaryl and substituted heteroaryl.

X₁ is selected from -OH, monoalkylamino, dialkylamino, alkoxide, arylkoxide and

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wherein:

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X₄ is hydroxide, arylamino, monoalkylamino, dialkylamino, alkoxide, or arylalkoxide;

R₃ is selected from the side chains of known natural α-amino acids, unnatural amino acids, hydrogen, 1-10 carbon linear and branched alkyl, 1-10 carbon linear and branched substituted alkyl, aryl, substituted aryl, 1-10 carbon linear and branched substituted aryl, alkoxyaryl, 3-8 carbon cycloalkyl, heterocycle, substituted heterocycle, heteroaryl and substituted heteroaryl.

2. The composition of claim 1 wherein X_i is

- 3. The composition of claim 2 wherein X_4 is -OH.
 - 4. The composition of claim 1 wherein X_4 is -OH.
- 5. The composition of claim 4 wherein R₁ is selected from 1-10 carbon branched alkyl, and 1-10 carbon unbranched alkyl substituents.
- The composition of claim 1 wherein X₄ is -OH, and R₁ and R₂ are each
 individually selected from side chains of the known natural α-amino acids, unnatural amino acids, and 1-10 carbon linear, alkyl and branched alkyl substituents.
 - 7. The composition of claim 6 wherein X₃ is selected from -C=O, -CH₂CO-, and (-CH₂)_a wherein n=0-2.
- 8. The composition of claim 7 wherein R₃ is selected from CO₂H, CH₂CO₂H, CH₂CO₂H, Arg, Lys, Asn, Gln, Asp, Glu, Phe and Nlc.

9. The composition of claim 8 wherein Ar is selected from indole and substituted indole.

- 10. The composition of claim 8 wherein Ar is selected from phenyl and substituted phenyl.
 - 11. The composition of claim 1 wherein X_2 is CH_2CO and R_1 is isobutyl.
- 10 12. The composition of claim 11 wherein X_2 is -OH, R_3 is H, X_3 , is H and Ar is selected from the group consisting of phenyl and indole.
 - 13. The composition of claim 11 wherein Ar is indole, R_1 is D-Leu (isobutyl), X_1 is H, and X_3 is -OH.
 - 14. The composition of claim 13 wherein R_2 is 2-NAP and R3 is Asp.
- 15. The composition of claim 13 wherein R_2 is 4,4'-BPA and R_3 is selected from the group consisting of Nle, Asp, Asn, β -Alanine, His, and Arg.
 - 16. The composition of claim 1 wherein Ar is indole, X_3 is selected from -C=O, and CH₂CO, R_3 is selected from biaryl and substituted biphenyl, R_1 is isobutane, R_3 is CH₂CO₂H and X_4 is -OH.
- 20 17. The composition of claim 1 wherein Ar is selected from phenyl and substituted phenyl, X₃ is selected from -C=O and -CH₂CO, R₂ is selected from biaryl and biphenyl, R₁ is isobutyl, R₃ is CH₂CO₂H, and X₄ is -OH.
 - 18. The composition of claim 1 wherein Ar is indole, X_3 is CH_2CO , R_2 is 4,4'-biphenyl, R_1 is isobutyl, R_3 is CH_2CO_2H , and X_4 is -OH.
- 25 19. A cationic salt of the composition of claim 1.
 - 20. An acid addition salt of the composition of claim 1.
 - 21. A method for inhibiting cancer in mammals comprising administering a therapeutically effective amount of the composition of claim 1 to the mammal.

22. The method of claim 21 wherein the therapeutically effective amount ranges from about 0.001 to about 100 mg/kg weight of the mammal.

- 23. The method of claim 21 wherein the composition is administered to a mammal suffering from auto immune disorders selected from the group consisting of lupus, MS, ARD and arthritits.
 - 24. The method of claim 23 wherein the disorder is RA.
 - 25. The method of claim 21 wherein the mammal is a human.
- 26. A pharmaceutical composition of matter comprising the composition of claim 1 and one or more pharmaceutical excipients.
- 27. The pharmaceutical composition of matter of claim 26 wherein the pharmaceutical composition is in the form of a solution.
 - 28. The pharmaceutical composition of matter of claim 26 wherein the pharmaceutical composition is in the form of a tablet.

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Inte onal Application No PCT/US 99/01097

			00 937 01037
A. CLASSII IPC 6	FICATION OF SUBJECT MATTER C07K5/03 C07K5/02 A61K38/04	3 C07K5/062	C07K5/083
According to	International Patent Classification (IPC) or to both national classifica	tion and IPC	
B. FIELDS	SEARCHED		
Minimum do IPC 6	cumentation searched (classification system followed by classification ${\tt C07K-A61K}$	n symbols)	
Documentat	ion searched other than minimum documentation to the extent that so	uch documents are included in the	ne fields searched
Electronic da	ata base consulted during the international search (name of data bas	e and, where practical, search to	erms used)
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT	,	
Category ^a	Citation of document, with indication, where appropriate, of the rele	evant passages	Relevant to claim No.
х	US 5 656 604 A (HEMMI KEIJI ET A 12 August 1997 The whole document; see especiall examples 79,81,153,155		1-10,19, 26-28
X	DA SETTIMO A ET AL: "Synthesis a benzodiazepine receptor affinity N-(indol-3-ylglyoxylyl)-dipeptide derivatives. Structural requireme inverse agonist/antagonist recept interactions" DRUG DES. DISCOVERY (DDDIEV, 10559612); 1993; VOL.10 (3 PP.199-211, XP002103282 Univ. Pisa; Ist. Chim. Farm.; Pisa Italy (IT) see the whole document	nts for the second or the seco	1,26-28
X Furt	her documents are listed in the continuation of box C.	X Patent family members	s are listed in annex.
"A" docume consider tilling of "L" docume which citatio "O" docume other "P" docume later t	ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international date ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another on or other special reason (as specified) lent referring to an oral disclosure, use, exhibition or means ent published prior to the international filing date but han the priority date claimed	cited to understand the prin invention "X" document of particular relevicannot be considered nove involve an inventive step w "Y" document of particular relevicannot be considered to involvement is combined with	onflict with the application but ciple or theory underlying the ance; the claimed invention of or cannot be considered to then the document is taken alone ance; the claimed invention wolve an inventive step when the tone or more other such docueing obvious to a person skilled time patent family
2	20 May 1999	04/06/1999	
Name and	mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Groenendijk	, M

Inte onal Application No
PCT/US 99/01097

C.(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category	Citation of document, with indication where appropriate, of the relevant passages	Relevant to claim No.
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X	CHEMICAL ABSTRACTS, vol. 089, no. 15, 9 October 1978 Columbus, Ohio, US; abstract no. 129909, LARSEN B R ET AL: "Products from the reaction of ninhydrin with triphenylalanine" XP002102781 see abstract & ANAL. BIOCHEM. (ANBCA2,00032697);1978; VOL.86 (1); PP.127-32, Univ. Texas Dent. Branch;Dep. Biochem.; Galveston; Tex.	
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Inte onal Application No
PCT/US 99/01097

	PCT/US 99/01097	
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category 3	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
, X	LUM R T ET AL: "Selective inhibition of the chymotrypsin-like activity of the 20S proteasome by 5-methoxy-1-indanone dipeptide benzamides" BIOORGANIC & MEDICINAL CHEMISTRY LETTERS, vol. 8, no. 3, 3 February 1998, page 209-214 XP004136850 see the whole document	1-7,11, 19-28
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rnational application No.

INTERNATIONAL SEARCH REPORT

PCT/US 99/01097

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Inte	rnational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely: Remark: Although claims 21-25 are directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.
2. X	Claims Nos.: $1-15$, $19-28$ (partial) because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
	see FURTHER INFORMATION sheet PCT/ISA/210
з. 🗀	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Inte	ernational Searching Authority found multiple inventions in this international application, as follows:
1.	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
З	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4.	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark	The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Claims Nos.: 1-15,19-28(all partially)

The scope of the claims 1-15 is very broad and speculative and/or said claims are ambigous and unclear. A formula consisting virtually of variables which are moreover in at least part of the claims ill-defined (e.g. the use of "aryl", "heteroaryls" and "sidechain of an unnatural amino acid) cannot be considered to be a clear and concise definition of patentable subject-matter (Art.6 PCT).

Furthermore the claims contain anumber of inconsistencies and ambigous defintions:

1)from the description it appears that the application relates to -ketoamides whereas claim 1 also encompasses compounds lacking said structural entity (compounds wherein X3 is (CH2)n);

2)in the claims 8, 14 and 15 (part of) the sidechains appear to consist of amino acid residues;

3)in claim 11 X2 has been defined as X3;

4)In the claims 12 and 13 the definitions of X2 and X3 are incompatible with claim 1.

Therefore a meaningful and economically feasible search could not encompass the complete subject-matter of the claims. Consequently the search had been directed to the claims 16-18 and the compounds defined in the examples (wherein X2 has been read as being X3) (closely) and also the claims 19-28 as far as relating to said compounds (Art.17(2)(a)(ii) and (b) PCT, PCT Guidelines CIII, 2.1 and CIII, 3,7).

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Inte onal Application No PCT/US 99/01097

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